# The phylo4 S4 classes and methods

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### March 7, 2008

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# 1 Introduction

This document describes the new phylo4 S4 classes and methods, which are intended to provide a unifying standard for the representation of phylogenetic trees and comparative data in R. The phylobase package was developed to help both end users and package developers by providing a common suite of tools

likely to be shared by all packages designed for phylogenetic analysis, facilities for data and tree manipulation, and standardization of formats.

For end-users, standardization will greatly simplify comparing analyses across different packages by easing data portability, as well as reducing the learning curve involved when using new packages. Users will also benefit by having a common repository of useful functions contained within one base package, for example tools for including or excluding subtrees (and associated phenotypic data) or improved tree and data plotting facilities. For developers, the phylobase package allows programming efforts to be put directly into developing new solutions for new problems (i.e. new phylogenetic methods) rather than re-coding the same base tools that each package requires. It is hoped that standardization will also synergize the efforts of individual developers into a comparative method community (this sounds stupid—please fix), as well as facilitating code validation by providing a repository for benchmark tests.

On a more abstract level, two motivations for the development of this package were better data checking and abstraction of the tree data formats. Currently phylobase is capable of checking that data and trees are associated in the proper fashion, and protects users and developers from accidently reordering one, but not the other. The phylobase package also seeks to abstract the data format so that commonly used information (for example, branch length information or the ancestor of a particular node) can be accessed without knowing the underlying data structure (i.e., whether the tree is stored as a matrix, or a list, or a parenthesis-based format). This is achieved through generic phylobase functions which which retrieve the relevant information from the data structures. The benefits of such abstraction are multiple: (1) easier access to the relevant information via a simple function call (this frees both users and developers from learning details of complex data structures), (2) freedom to optimize data structures in the future without breaking code. Having the generic functions in place to "translate" between the data structures and the rest of the program code allows program and data structure development to proceed somewhat independently. The alternative is code written for specific data structures, in which modifications to the data structure requires rewriting the entire package code (often exacting too high a price, which results in the persistence of less-optimal data structures). (3) providing broader access to the range of tools in phylobase. Developers of specific packages can use these new tools based on S4 objects without knowing the details of S4 programming.

The base phylo4 class is modeled on the the phylo class in ape. phylo4d and multiphylo4 extend the phylo4 class to include data or multiple trees respectively. In addition to describing the classes and methods this vignette gives examples of how they might be used.

# 2 Package Overview

The phylobase package currently implements the following functions and data structures:

- Data structures for storing a single tree and multiple trees: phylo4 and multiPhylo4?
- A data structure for storing a tree with associated tip and node data: phylo4d
- A data structure for storing multiple trees with one set of tip data: multiPhylo4d
- Functions for reading nexus files into the above data structures
- Functions for converting between the above data structures and ape phylo objects as well as ade4 phylog objects
- Functions for editing trees and data (i.e., subsetting and replacing)
- Functions for plotting trees and trees with data

# 3 Using the S4 help system

The S4 help system works similarly to the S3 help system with some small differences relating to how S4 methods are written. The plot() function is a good example. When we type ?plot we are provided the help for the default plotting function which expects x and y. R also provides a way to smartly dispatch the right type of plotting function. In the case of an ape phylo object (a S3 class object) R evaluates the class of the object and finds the correct functions, so the following works correctly.

```
> library(ape)
> rand_tree <- rcoal(10)
> plot(rand_tree)
```

However, typing ?plot still takes us to the default plot help. We have to type plot.phylo to find what we are looking for. This is because S3 generics are simply functions with a dot and the class name added.

The S4 generic system is too complicated to describe here, but doesn't include the same dot notation. As a result <code>?plot.phylo4</code> doesn't work, R does, however, find the right plotting function.

```
> library(phylobase)
> rand_p4_tree <- as(rand_tree, "phylo4")
> plot(rand_p4_tree)
```

All fine and good, but how to we find out about all the great features of the phylobase plotting function? R has two nifty ways to find it, the first is to simply put a question mark in front of the whole call:

```
> ?plot(rand_p4_tree)
```

R looks at the class of the rand\_p4\_tree object and takes us to the correct help file (note: this only works with S4 objects). The second ways is handy if you already know the class of your object, or want to compare to generics for different classes:

```
> method?plot("phylo4")
```

More information about how \$4 documentation works can be found in the methods package, by running the following command.

```
> help("Documentation", package = "methods")
```

### 4 Trees without data

You can start with a tree — an object of class phylo from the ape package (e.g., read in using the read.tree() or read.nexus() functions), and convert it to a phylo4 object.

For example, load the raw Geospiza data:

```
> data(geospiza_raw)
> names(geospiza_raw)
```

```
[1] "tree" "data"
```

Convert the S3 tree to a S4 phylo4 object using the as() function:

> library(phylobase)
> g1 <- as(geospiza\_raw\$tree, "phylo4")
> g1

	taxon.name	node	ancestor	branch.length	node.type
1	NO1	15	NA	NA	root
2	NO2	16	15	0.29744	internal
3	NO3	17	16	0.04924	internal
4	NO4	18	17	0.06859	internal
5	N05	19	18	0.13404	internal
6	N06	20	19	0.10346	internal
7	NO7	21	20	0.03550	internal
8	N08	22	21	0.00917	internal
9	N09	23	22	0.07333	internal
10	N10	24	23	0.05500	internal
11	N11	25	19	0.24479	internal
12	N12	26	25	0.05167	internal
13	N13	27	26	0.01500	internal
14	fuliginosa	1	24	0.05500	tip
15	fortis	2	24	0.05500	tip
16	${\tt magnirostris}$	3	23	0.11000	tip

tip
tip

Note that the nodes and edges are given default names if the tree contains no node or edge names.

The summary method gives a little extra information, including information on branch lengths:

### > summary(g1)

No root edge.

Phylogenetic tree: g1

Number of tips : 14 Number of nodes : 13

Branch lengths:

mean : 0.1764008 variance : 0.04624379

distribution :

Min. 1st Qu. Median 3rd Qu. Max. 0.00917 0.04985 0.08000 0.21910 0.88080

Print tip labels:

### > labels(g1)

```
[1] "fuliginosa" "fortis" "magnirostris" "conirostris" "scandens" [6] "difficilis" "pallida" "parvulus" "psittacula" "pauper" [11] "Platyspiza" "fusca" "Pinaroloxias" "olivacea"
```

Print internal node labels (R automatically assigns values):

#### > nodeLabels(g1)

```
[1] "N01" "N02" "N03" "N04" "N05" "N06" "N07" "N08" "N09" "N10" "N11" "N12" [13] "N13"
```

Print edge labels (also automatically assigned):

```
> edgeLabels(g1)
 [1] "E16" "E17" "E18" "E19" "E20" "E21" "E22" "E23" "E24" "E1" "E2" "E3"
[13] "E4" "E5" "E6" "E25" "E7" "E26" "E27" "E8" "E9" "E10" "E11" "E12"
[25] "E13" "E14"
  Is it rooted?
> isRooted(g1)
[1] TRUE
   Which node is the root?
> rootNode(g1)
[1] 15
  Does it have any polytomies?
> hasPoly(g1)
[1] FALSE
  Does it have branch lengths?
> hasEdgeLength(g1)
[1] TRUE
   You can modify labels and other aspects of the tree — for example,
> labels(g1) <- tolower(labels(g1))</pre>
```

### 5 Trees with data

The phylo4d class matches trees with data. (fixme: need to be able to use ioNCL!) or combine it with a data frame to make a phylo4d (tree-with-data) object.

Now we'll take the *Geospiza* data from geospiza\_raw\$data and merge it with the tree. However, since *G. olivacea* is included in the tree but not in the data set, we will initially run into some trouble:

```
> g2 <- phylo4d(g1, geospiza_raw$data)
    gives

Error in check_data(res, ...) :
    Tip data names are a subset of tree tip labels
(missing data names: platyspiza,pinaroloxias,olivacea)
(extra data names: Platyspiza,Pinaroloxias)</pre>
```

We have two problems — the first is that we forgot to lowercase the labels on the data to match the tip labels:

```
> gdata <- geospiza_raw$data
> row.names(gdata) <- tolower(row.names(gdata))</pre>
```

To deal with the second problem (missing data for *G. olivacea*), we have a few choices. The easiest is to use missing.tip.data="OK" to allow R to create the new object:

```
> g2 <- phylo4d(g1, gdata, missing.tip.data = "OK")</pre>
```

(setting missing.tip.data to "warn" would create the new object but print a warning).

Another way to deal with this would be to use prune() to drop the offending tip from the tree first:

```
> g1B <- prune(g1, "olivacea")
> phylo4d(g1B, gdata)
```

You can summarize the new object:

> summary(g2)

No root edge.

Phylogenetic tree : as(object, "phylo4")

Number of tips : 14 Number of nodes : 13

Branch lengths:

mean : 0.1764008 variance : 0.04624379

 ${\tt distribution} \ :$ 

Min. 1st Qu. Median 3rd Qu. Max. 0.00917 0.04985 0.08000 0.21910 0.88080

#### Comparative data:

Tips: data.frame with 14 taxa and 5 variables

wingL		tarsusL		culmenL		beakD	
Min.	:3.975	Min.	:2.807	Min.	:1.974	Min.	:1.191
1st Qu.	:4.189	1st Qu.	:2.929	1st Qu.	:2.187	1st Qu	:1.941
Median	:4.235	Median	:2.980	Median	:2.311	${\tt Median}$	:2.073
Mean	:4.236	Mean	:2.991	Mean	:2.333	Mean	:2.083
3rd Qu.	:4.265	3rd Qu.	:3.039	3rd Qu.	:2.430	3rd Qu	:2.347
Max.	:4.420	Max.	:3.271	Max.	:2.725	Max.	:2.824

```
NA's
       :1.000
                NA's
                       :1.000
                                 NA's
                                        :1.000
                                                  NA's
                                                          :1.000
    gonysW
Min.
       :1.401
1st Qu.:1.845
Median :1.962
Mean
       :2.014
3rd Qu.:2.222
       :2.676
Max.
NA's
       :1.000
```

Object contains no node data.

Or use tdata() to extract the data (i.e., tdata(g2)). By default, tdata() will retrieve tip data, but you can also get internal node data only (tdata(tree, "node")) or — if the tip and node data have the same format — all the data combined (tdata(tree, "allnode")).

Plotting calls plot.phylog from the ade4 package.

If you want to plot the data (e.g. for checking the input), plot(tdata(g2)) will create the default plot for the data — in this case, since it is a data frame [this may change in future versions but should remain transparent] this will be a pairs plot of the data.

## 6 Subsetting

The subset command offers a variety of ways of extracting portions of a phylo4 or phylo4d tree, keeping any tip/node data consistent.

tips.include give a vector of tips (names or numbers) to retain

tips.exclude give a vector of tips (names or numbers) to drop

**mrca** give a vector of node or tip names or numbers; extract the clade containing these taxa

**node.subtree** give a node (name or number); extract the subtree starting from this node

Different ways to extract the fuliginosa-scandens clade:

```
> subset(g2, tips.include = c("fuliginosa", "fortis", "magnirostris",
+    "conirostris", "scandens"))
> subset(g2, node.subtree = "NO7")
> subset(g2, mrca = c("scandens", "fortis"))
    One could drop the clade by doing
> subset(g2, tips.exclude = c("fuliginosa", "fortis", "magnirostris",
+    "conirostris", "scandens"))
> subset(g2, tips.exclude = names(descendants(g2, MRCA(g2, c("difficilis",
+    "fortis")))))
```

Another approach is to pick the subtree graphically, by plotting the tree and using identify, which returns the identify of the node you click on with the mouse.

```
> plot(g1)
> n1 <- identify(g1)
> subset(g2, node.subtree = n1)
```

## 7 Tree-walking

getnodes, children, parent, descendants, ancestors, siblings, MRCA... generally take a phylo4 object, a node (specified by number or name) and return a named vector of node numbers.

# 8 multiPhylo classes

## 9 Examples

# A Definitions/slots

### A.1 phylo4

Like phylo, the main components of the phylo4 class are:

edge an  $N \times 2$  matrix of integers, where the first column ...

edge.length numeric list of edge lengths (length N or empty)

Nnode integer, number of nodes

tip.label character vector of tip labels (required)

node.label character vector of node labels (maybe empty)

root.edge integer defining root edge (maybe NA)

We have defined basic methods for phylo4:show, print (copied from print.phylo inape), and a variety of accessor functions (see help files). summary does not seem to be terribly useful in the context of a "raw" tree, because there is not much to compute: end users?

Print method: add information about (ultrametric, scaled, polytomies (zero-length or structural))?

### A.2 phylo4d

The phylo4d class extends phylo4 with data. Tip data, (internal) node data, and edge data are stored separately, but can be retrieved together or separately with tdata(x,"tip") or tdata(x,"all").

edge data can also be included — is this useful/worth keeping?

### A.3 multiphylo4

# B Validity checking

- number of rows of edge matrix (N) == length of edge-length vector (if > 0)
- (number of tip labels)+(nNode)-1 == N
- data matrix must have row names
- row names must match tip labels (if not, spit out mismatches)

•

Default node labels:

# C Hacks/backward compatibility

There is a way to hack the \$ operator so that it would provide backward compatibility with code that is extracting internal elements of a phylo4. The basic recipe is:

```
> setMethod("$", "phylo4", function(x, name) {
+    attr(x, name)
+ })
```

but this has to be hacked slightly to intercept calls to elements that might be missing. For example, ape detects whether log-likelihood, root edges, node labels, etc. are missing by testing whether they are NULL, whereas missing items are represented in phylo4 by zero-length vectors in the slots (or NA for the root edge) — so we need code like

```
> if (!hasNodeLabels(x)) NULL else x@node.label
```

to handle these cases.